

**SINTERED MATERIAL PRODUCT AND METHOD FOR
MANUFACTURING THE SAME**

BACKGROUND OF THE INVENTION

5 The present invention relates to a sintered material
product used for transportation machines such as
automobiles, ships, other general industrial machinery,
household electric appliances, OA equipment, and building
materials or printing. Particularly, the present invention
10 relates to a sintered material product having good sliding
properties and good corrosion resistance.

Copper alloys, which have good heat, electrical
conduction and good corrosion resistance, and good
workability, have been used in various industrial fields
15 such as buildings, printings, electrical facilities,
automobiles, and others. Particularly, copper alloys with
good corrosion resistance and good sliding properties are
broadly used for sliding parts in various gears, guide
rails, automobile parts, household electric appliances,
20 pumps, motors, OA equipment, and others.

In environments where the lubricating oil cannot be
used and a long-term durability is required, the slide
surface is subjected to a surface treatment;

For example, it has been proposed to form on the slide
25 surface a fluorocarbon resin coating, which is superior in

weathering resistance, dirt prevention, water repellent,
and lubricating characteristic. Alternatively, it has been
proposed to form on the slide surface a functional ceramic
film, which has heat resistance, wear resistance,
5 hydrophilicity, photocatalyst property, and far-infrared
reflection function.

However, the above-mentioned films generally have poor
adhesion to a sintered copper alloy being a substrate,
thus resulting in poor durability.

10 JP-P1993-157115A discloses the method of impregnating
a resin, such as polyamide or polyimide, into a sintered
copper alloy. The above-mentioned resins tend to be easily
impregnated into pores of the sintered copper alloy, thus
providing good adhesion to the sintered alloy material.

15 However, compared with films of fluorocarbon resin
such as PTFE (polytetrafluoro-ethylene), the above-
mentioned resin films are poor in sliding properties, so
that a sufficient sliding performance cannot be obtained.

In contrast, compared with resins such as polyamide
20 and polyimide, it is difficult to impregnate fluorocarbon
resins such as PTFE into pores of a sintered copper alloy.
Hence, the fluorocarbon resins show poor adhesive property
to copper alloys.

A solid lubricant such as molybdenum disulfide
25 indicates a tendency similar to those of fluorocarbon

resins.

In order to improve the film adhesive property, it has been considered to blast the sintered copper alloy with sands or ceramic particles.

5 This blasting technique improves the adhesive property but blast particles pierce the substrate (of a soft copper alloy). The pierced blast particles wear and deform the opposite material and degrade the seizure resistance or durability. For that reason, the blasting technique cannot
10 be employed to sintered copper alloy products.

SUMMARY OF THE INVENTION

The present invention is made to solve the above-mentioned problems.

15 An object of the present invention is to provide a sintered copper alloy material product having good sliding properties and good durability.

 Another object of the present invention is to provide a sintered copper alloy material product on which a skin
20 (particularly, a skin of a solid lubricant with low adhesion, such as PTFE or molybdenum disulfide) is securely formed on the surface thereof.

 Further another object of the present invention is to provide a sintered copper alloy material product having
25 good sliding properties and good durability under

unlubricating environments or corrosive environments.
The present inventor has studied aggressively to solve the above-mentioned problems. As a result, a component prone to corrosion (an easy-corrosive component) such as zinc contained in a copper alloy corrodes under severe

5 corrosive environment so that an oxidation product is produced. It was found that the corrosion accelerates seizure. In order to prevent the earlier seizure, it was considered to apply a solid lubricant, such as PTFE or molybdenum disulfide, on the surface of the substrate (a sintered copper alloy material). However, the solid
10 lubricant shows poor adhesion to the substrate (a copper alloy material). Consequently, this approach proved that a sufficient effect could not be obtained.

15 Further investigation has been aggressively made by the present inventor. Using a solution containing a compound of one kind or more selected from the group consisting of peroxide, peroxocompound, chromic acid and permanganic acid, easy-corrosive components or oxide
20 products were removed selectively. Thereafter, a skin of a solid lubricant such as PTFE or molybdenum disulfide was formed. Products produced thus have the skin invaded into pores of the substrate (a sintered copper alloy material). This conformation has an improved skin adhesion.
25 Furthermore, it has been known that the removal of the

corrosive product leads to an improved durability (corrosion resistance).

The present invention was made based on the above-mentioned knowledge.

5 The above-mentioned problems are solved by the following manufacturing methods. That is, a method for manufacturing a sintered material product having a sintered copper alloy material and a skin formed on the sintered copper alloy material, comprises the steps of (A)
10 processing said sintered copper alloy material with a selective chemical etching solution, and (C) forming a skin on the surface of said sintered copper alloy material after the step (A).

 Particularly, a method for manufacturing a sintered
15 material product having a sintered copper alloy material and a skin formed on the sintered copper alloy material, comprises the step of (A) processing the sintered copper alloy material with a selective chemical etching solution, (B) forming a layer of a metal phosphate and/or a metal
20 oxide on the surface of the sintered copper alloy material after the step (A), and (C) forming a skin on the surface of the sintered copper alloy material after the step (B).

 The processing temperature and the processing time of selective chemical etching solution are not restricted,
25 unless specified. The processing temperature is, for

example, a room temperature to 50° C. The processing time is, for example, 1 to 10 minutes.

The sintered product produced thus has the following structure.

5 That is, the sintered copper alloy material has pores of which the inlet diameter ranges from 10 μ m to 200 μ m. The average value of (inlet pore diameter)/(inner pore diameter) of each pore is 2 or more. Particularly, (inlet pore diameter)/(inner pore diameter) is 2 to 20. The
10 porosity is 2 to 35 % by volume.

Moreover, a layer of a metal phosphate and/or metal oxide is formed underneath of the skin.

The selective chemical etching solution used in the step (A) is, preferably, a solution containing a compound
15 of one kind or more selected from the group consisting of peroxide, peroxocompound, chromic acid, and permanganic acid. Particularly, the etching solution is, preferably, an aqueous solution containing peroxide or peroxocompound.

As peroxocompound are listed, for example,
20 peroxosulfuric acid, peroxophosphoric acid, peroxovanadic acid, peroxoniobic acid, peroxotantallic acid, peroxoboric acid, peroxotitanic acid, peroxotungstic acid, peroxomolybdic acid, and peroxochromic acid. Soluble salt of the above-mentioned acids, particularly, peroxosulfuric
25 acid and salt thereof, may be used. Particularly, ammonium

salt, sodium salt, and potassium salt of peroxodisulfuric acid is preferable.

The peroxide is preferably hydrogen peroxide.

The preferable concentration of peroxide or
5 peroxocompound is 1 to 30 % by weight (particularly, 3 to 20 % by weight). The reason is that an excessively low concentration results in a small selective removing effect of easy-corrosive components or oxides. In contrast, an excessively high concentration results in precipitation of
10 crystals and instability of etching solution.

The selective chemical etching solution contains, preferably, peroxide or peroxocompound, as well as a compound of one kind or more selected from the group consisting of phosphoric acid, sulfuric acid, nitric acid,
15 hydrochloric acid, hydrofluoric acid, zirconic hydrofluoric acid, titanitic hydrofluoric acid, titanitic acid, molybdic acid, tungstic acid, vanadic acid, niobic acid, and organic chelating agent (preferably, tartaric acid, citric acid, EDTA, organic phosphonic acid, or phytic acid,
20 having a chelating effect to copper).

A selective chemical etching solution, having a preferable combination, is an aqueous solution including, for example, phosphoric acid-peroxosulfuric acid, peroxophosphoric acid-sulfuric acid, phosphoric acid-
25 hydrogen peroxide, tartaric acid-hydrogen peroxide,

phosphoric acid-nitric acid-hydrogen peroxide, or sulfuric acid-hydrogen peroxide.

pH of the selective chemical etching solution is preferably 1 to 5. pH is adjusted appropriately using
5 alkalis such as aqueous ammonia, ammonium carbonate, sodium hydroxide, sodium carbonate, or potassium hydroxide.

In the present invention, peroxide and peroxocompound are especially used for the following reasons. That is, the desired selective etching cannot be realized though
10 the mechanical surface roughening method using the blasting process or through the chemical etching using common acid or alkali. The selective etching can be effectively realized using peroxide or peroxocompound. The skin is formed on the sintered copper alloy material after
15 the step (A) so as to invade into the surface thereof. This can provide a very high anchor effect and high skin adhesion.

After the step (A), it is preferable to process the sintered copper alloy material using a solution containing
20 a metal compound of one kind or more selected the group consisting of Zn, Ca, Mg, Mn, Ni, Co, Mo, W, Cu, Sn, Ti, Zr, V, In, and Cr.

For example, a sintered copper alloy material is processed using a metal ion aqueous solution of sulfate,
25 acetate, chloride, phosphate, carbonate, or hydroxide of

the above-mentioned metals or using colloidal sol of the above-mentioned metals oxide. Thus, a metal phosphate layer and/or a metal oxide layer is formed on the surface of the sintered copper alloy material subjected to the selective chemical etching. A preferable thickness of the layer is 0.1 to 2 μm (preferably, 0.2 to 1 μm). This layer improves the corrosion resistance. The adhesiveness between the layer and the skin formed on the layer is improved.

After the step (A) and the step (B), it is preferable to rinse quickly the sintered copper alloy material in water. In other words, acids left in the sintered copper alloy material are removed by water washing. For example, the acid remaining in the sintered copper alloy material causes a degradation of corrosion resistance. It is particularly preferable to subject the sintered copper alloy material to ultrasonic cleaning. That is, the smut remaining in the surface of the sintered copper alloy material is removed through the ultrasonic cleaning, so that the material surface with good adhesive property can be obtained.

After the water rinsing, it is preferable to process the sintered copper alloy material with an aqueous solution containing organic alkali compound (e.g. alkanolamine). This process neutralizes the acid remaining

in the sintered copper alloy material, thus improving the corrosion resistance. Moreover, the adhesive property of the skin formed on the sintered copper alloy material is improved. Low molecular compounds, each of which the molecular structure has at least one amino group, are preferable as organic alkali compounds. For example, as low molecular compounds are listed monoethanolamine, diethanolamine, triethanolamine, morpholine, derivatives of them, and alkoxysilane having various amino groups. These compounds improve the surface adhesive property of a solid lubricative skin and the wetting property of paint.

A skin (particularly, a solid lubricant skin) is formed on the surface of the sintered copper alloy material processed as shown above.

A coating film containing a solid lubricant is listed as the skin. As solid lubricants are appropriately used, for example, polytetrafluoro-ethylene (PTFE), tetrafluoroperfluoroalkylvinylethercopolymer (PFA), graphite, molybdenum disulfide, tungsten disulfide, boron nitride, tungsten fluoride, titanium nitride. As binder components of a coating film are listed, for example, resins such as polyester, polyolefin, polyurethane, polyacryl, polyamide, polyimide, epoxy, and silicone. As these paints can be used, for example, FL-J4668 produced by Nihon Parkerizing Co., Ltd. and EB3 and LHF4B produced

by Kawamura Research Laboratories, Inc. A paint containing the various components is applied on the sintered copper alloy material using the splay method, the dipping method, the roll coating method, the powder method or the electro-
5 deposition method and then is baked, so that a painted film is formed. The thickness of the painted film is preferably, for example, 1 to 40 μm (particularly, the film thickness is 2 to 20 μm or 1 to 10 μm).

According to the present invention, the sintered
10 copper alloy contains copper. The content of copper is preferably 20 to 95 % by weight (is, particularly, 50 to 90 % by weight). In addition, as a component other than Cu is listed, for example, Zn, Fe, Mn, Al, Co, P, Mo, Sn, Ni or C. Specifically, the sintered copper alloy material is
15 B031, B110, B062, or B060.

Prior to the selective chemical etching process, it is preferable to degrease and clean the sintered copper alloy material. Thus, oil contents adhering to the surface of the sintered copper alloy material are removed.

20 The products produced as described above proved that the durability and the seizure resistance (sliding property) are remarkably improved.

DESCRIPTION OF THE EMBODIMENTS

25 A sintered material product according to the present

invention includes a sintered copper alloy material and a skin formed on the sintered copper alloy material. The skin is, particularly, a paint film containing a solid lubricant.

5 The sintered copper alloy material has pores. The inlet of each pore has a diameter of 10 to 200 μm (particularly, 20 to 100 μm). An average value of (inlet pore diameter)/(inner pore diameter) is 2 or more (particularly, 2 to 20, or 5 to 20). The porosity is 2 to 10 35 % by volume (particularly, 10 to 25 % by volume).

 The porous sintered copper alloy material is obtained by processing a sintered copper alloy material with a selective chemical etching solution. For example, a sintered copper alloy material is dipped in a selective 15 chemical etching solution. Alternatively, a selective chemical etching solution may be sprayed onto the sintered copper alloy material. In other way, the surface of the sintered copper alloy material may be wetted with a selective chemical etching solution according to the flow 20 coating method. The processing temperature is a room temperature to 50 ° C. The processing time is about 1 to 10 minutes.

 In the sintered material product of the present invention, a layer of metal phosphate and/or metal oxide 25 is formed between the skin and the sintered copper alloy

material.

The present invention relates to the method for manufacturing a sintered material product having a sintered copper alloy material and a skin formed on the sintered copper alloy material. This manufacturing method includes the step (A) of processing a sintered copper alloy material with a selective chemical etching solution. After the step (A), the manufacturing method has the step (C) of forming a skin on the surface of the sintered copper alloy material. The manufacturing method further has the step (B) of forming a layer of metal phosphate and/or metal oxide on the surface of the sintered copper alloy material between step (A) and the step (C).

The selective chemical etching solution used in the step (A) is preferably a solution containing a compound of one kind or more selected from the group consisting of peroxide, peroxocompound, chromic acid and permanganic acid. Particularly, the etching solution is preferably an aqueous solution containing peroxide or peroxocompound.

As peroxocompounds are listed peroxosulfuric acid, peroxophosphoric acid, peroxovanadic acid, peroxoniobic acid, peroxotantallic acid, peroxoboric acid, peroxotitanic acid, peroxotungstic acid, peroxomolybdic acid, and peroxochromic acid. Soluble salts of the above-mentioned acids may be used. Particularly, peroxosulfuric acid or

salt thereof, may be used. Particularly, ammonium salt, sodium salt, and potassium salt of peroxodisulfuric acid is preferable. The peroxide is preferably hydrogen peroxide.

5 The preferable concentration of peroxide or peroxocompound is 1 to 30 % by weight (particularly, 3 to 20 % by weight).

 The selective chemical etching solution contains, preferably, peroxide or peroxocompound, as well as a
10 compound of one kind or more selected from the group consisting of phosphoric acid, sulfuric acid, nitric acid, hydrochloric acid, hydrofluoric acid, zirconic hydrofluoric acid, titanitic hydrofluoric acid, titanitic acid, molybdic acid, tungstic acid, vanadic acid, niobic acid,
15 and organic chelating agent (for example, tartaric acid, citric acid, EDTA, organic phosphonic acid, or phytic acid).

 A selective chemical etching solution, having a preferable combination, is an aqueous solution including,
20 for example, phosphoric acid-peroxosulfuric acid, peroxophosphoric acid-sulfuric acid, phosphoric acid-hydrogen peroxide, tartaric acid-hydrogen peroxide, phosphoric acid-nitric acid-hydrogen peroxide, or sulfuric acid-hydrogen peroxide.

25 pH of the selective chemical etching solution is

preferably 1 to 5. pH is adjusted appropriately using alkalis such as aqueous ammonia, ammonium carbonate, sodium hydroxide, sodium carbonate, or potassium hydroxide.

5 If the selective chemical etching solution does not contain acid (e.g. phosphoric acid) or organic chelating agent, it is considered that after the selective chemical etching process, post treatment may be carried out with acid (e.g. phosphoric acid) or with an organic chelating agent. However, this approach results in an increased
10 number of production steps.

After the step (A), the step (B) is carried out, if necessary. In other words, after the selective chemical etching process, the sintered copper alloy material is processed with a solution containing metal compounds. A
15 preferable metal, used in such a step, is Zn, Ca, Mg, Mn, Ni, Co, Mo, W, Cu, Sn, Ti, Zr, V, In, or Cr. For example, a sintered copper alloy material is processed using an aqueous solution of sulfate, acetate, chloride, phosphate, carbonate, or hydroxide of the above-mentioned metals
20 (having a concentration of 0.02 to 2 % by weight). Alternatively, the sintered copper alloy material may be processed using colloidal sol of a metal oxide. Thus, a layer of metal phosphate and/or metal oxide, having a thickness of 0.1 to 2 μm (preferably, 0.2 to 1 μm), is
25 formed.

After the step (A) or the step (B), it is preferable, if necessary, to rinse quickly the sintered copper alloy material in water (particularly, to perform ultrasonic cleaning to it). This process removes acids left in the sintered copper alloy material.

After the water rinsing, the sintered copper alloy material is processed, if necessary, with an aqueous solution containing organic alkali compound. Low molecular compounds, each of which the molecular structure has at least one amino group, are listed as organic alkali compounds. For example, as low molecular compounds are listed monoethanolamine, diethanolamine, triethanolamine, morpholine, derivatives of them, and alkoxysilane having various amino groups.

After the above-mentioned process, a skin is formed on the surface of the sintered copper alloy material. Particularly, the film is a paint (or coating) film containing a solid lubricant. As solid lubricants are listed, for example, PTFE, TPA, graphite, molybdenum disulfide, tungsten disulfide, boron nitride, tungsten fluoride, titanium nitride. As paint film binder components are listed, for example, resins (the thermoplastic resins) such as polyester, polyolefin, polyurethane, polyacryl, polyamide, polyimide, epoxy, and silicone. As these paints can be used, for example, FL-

J4668 produced by Nihon Parkerizing Co., Ltd. and EB3 and LHF4B produced by Kawamura Research Laboratories, Inc. A paint containing the various components mentioned above is applied on the sintered copper alloy material using the splay method, the dipping method, the roll coating method, the powder method or the electro-deposition method and then is baked. Thus, a painted film having, for example, a thickness of 2 to 20 μm (particularly, 1 to 10 μm) is formed.

According to the present invention, the sintered copper alloy is one containing copper. The content of copper is preferably 20 to 95 % by weight (is, particularly, 50 to 90 % by weight). In addition, a component, other than Cu is, for example, Zn, Fe, Mn, Al, Co, P, Mo, Sn, Ni or C. Specifically, the sintered copper alloy material is B031, B110, B062, or B060.

Specific embodiments will be described below together with a comparative example.

[Embodiment 1]

Copper alloy powder, which has a grain size of 150 μm or less, was molded in a predetermined shape under a molding pressure of 150 to 350 MPa. Thereafter, the molded material was sintered for 40 minutes in an atmosphere of an ammonium decomposition gas at 700 to 900 ° C. The sintered material was corrected with a sizing pressure of

100 to 400 MPa.

First, the sliding parts made of the resultant sintered copper alloy material B031 (Cu of 85% by weight, Sn of 10% by weight and C of 5% by weight) was degreased. The degreasing agent was alkali degreasing agent (FC-315 produced by Nihon Parkerizing Co., Ltd.). The concentration of the degreasing agent was 20 g/l. The degreasing temperature was 60 ° C. The degreasing time was 2 minutes.

After degreasing, the sliding parts were rinsed in water. Thereafter, the sliding parts were dipped in the aqueous solution containing peroxodisulfate potassium (of a concentration of 5% by weight) and sulfuric acid (of a concentration of 5% by weight). The dipping temperature was 27 ° C. The dipping time was 180 seconds.

After dipping, the sliding parts were cleaned in running water.

Thereafter, FL-J4668 (containing a main constituent of PTFE) (produced by Nihon Parkerizing Co., Ltd) was sprayed with a spray gun so as to have a dry thickness of 3 μ m. Then, the sprayed coating was baked at 200 ° C.

[Embodiment 2]

B031-made sliding parts, obtained in a manner similar to that of the embodiment 1, were used.

First, the sliding parts were degreased. The

degreasing agent was FC-315. The concentration of the degreasing agent was 20 g/l. The degreasing temperature was 60 ° C. The degreasing time was 2 minutes.

5 After degreasing, the sliding parts were rinsed in water.

Thereafter, the sliding parts were dipped in the aqueous solution containing peroxophosphoric sodium (of a concentration of 5% by weight) and phosphoric acid (of a concentration of 10% by weight). The dipping temperature
10 was 27 ° C. The dipping time was 180 seconds.

After dipping, the sliding parts were cleaned in running water.

Thereafter, LHF4B (containing a main constituent of PTFE) (produced by Kawamura Research Laboratories, Inc.)
15 was sprayed with a spray gun so as to have a dry thickness of 4 μm. Then, the sprayed coating was baked at 200 ° C.
[Embodiment 3]

B031-made sliding parts, obtained in a manner similar to that of the embodiment 1, were used.

20 First, the sliding parts were degreased. The degreasing agent, FC-315, was used. The concentration of the degreasing agent was 20 g/l. The degreasing temperature was 60 ° C. The degreasing time was 2 minutes.

25 After degreasing, the sliding parts were rinsed in water.

Thereafter, the sliding parts were dipped in the aqueous solution containing chromic acid (of a concentration of 5% by weight) and hydrogen peroxide (of a concentration of 5% by weight). The dipping temperature was 27 ° C. The dipping time was 180 seconds.

After dipping, the sliding parts were cleaned in running water.

Next, the sliding parts were dipped in an aqueous solution containing phosphoric acid (of 5% by weight) and zinc nitrate (2% by weight). The dipping temperature was 45 ° C. The dipping time was 60 seconds.

Then, the sliding parts were subjected to ultrasonic cleaning (25 KHz x 30 minutes).

Thereafter, FL-J4668 was sprayed with a spray gun so as to form a dry thickness of 3 μ m. Then, the sprayed coating was baked at 200 ° C.

[Embodiment 4]

B031-made sliding parts, obtained in a manner similar to that of the embodiment 1, were used.

First, the sliding parts were degreased. The degreasing agent, FC-315, was used. The concentration of the degreasing agent was 20 g/l. The degreasing temperature was 60 ° C. The degreasing time was 2 minutes.

After degreasing, the sliding parts were rinsed in water.

Thereafter, the sliding parts were dipped in an aqueous solution containing peroxodisulfate potassium (of a concentration of 5% by weight) and sulfuric acid (of a concentration of 5% by weight). The dipping temperature was 27 ° C. The dipping time was 180 seconds.

After dipping, the sliding parts were cleaned in running water.

Next, the sliding parts were dipped in an aqueous solution of monoethanolamine (of a concentration of 0.5% by weight) and then were lift up after a lapse of 15 seconds. Successively, the sliding parts were dried at 120 ° C for 10 seconds.

Thereafter, FL-J4668 was sprayed with a spray gun so as to form the dry thickness of 3 μ m. Then, the sprayed coating was baked at 200 ° C.

[Comparative Example 1]

B031-made sliding parts, obtained in a manner similar to that of the embodiment 1, were used.

First, the sliding parts were degreased. The degreasing agent was FC-315. The concentration of the degreasing agent was 20 g/l. The degreasing temperature was 60 ° C. The degreasing time was 2 minutes.

After degreasing, the sliding parts were rinsed in water.

Thereafter, the sliding parts were dipped in the

aqueous solution containing hydrochloric acid (of a concentration of 10% by weight). The dipping temperature was 27 ° C. The dipping time was 180 seconds.

After dipping, the sliding parts were cleaned in running water.

Thereafter, FL-J4668 was sprayed with a spray gun so as to form a dry thickness of 3 μm. Then, the sprayed coating was baked at 200 ° C.

[Characteristics]

The sliding parts obtained by the above-mentioned embodiments were inspected in adhesive property, corrosion resistance and sliding property of a paint film (a lubricating film). Table 1 shows the results. Moreover, the surface profiles of the B031 materials after etching process were inspected. Table 1 shows the results.

Table 1

		Paint Film			B031 Material Surface	
		Adhesive property	Corrosion resistance	Sliding property	Porosity	Pore size ratio
20	Embodiment 1	○	○	○	27.9	15.6
	Embodiment 2	○	○	○	30.6	16.7
	Embodiment 3	○	○	○	25.1	13.4
	Embodiment 4	◎	○	◎	33.5	18.9
	Comparative Ex. 1	×	△	×	1.4	1.2

Adhesive property (cross-cut test):

A paint film was cut with a cutter knife to form 100 gridirons, each having 1 mm x 1 mm. Next, an adhesive cellophane tape is applied onto the paint film and then peeled off. In this test, symbol ◎ represents the case where no skin is peeled off. Symbol ○ represents the case where one or three pieces of skin are peeled off. Symbol △ represents the case where 4 or 20 pieces of skin are peeled off. Symbol × represents the case where 21 pieces or more of skin are peeled off.

Corrosion resistance:

A salt splay test was carried out in conformity with JIS-K5400. Symbol ○ indicates that there is no sign of generation of rust. Symbol △ indicates that there is a sign of generation of rust. In the case where rust is recognized, the material is handled as a rejected article.

Sliding property:

The load (N) until seizure occurs was inspected using the SRV friction/wear tester. The load to be applied was stepped up at a rate of 50 N/min. The frequency of vibration is 50 Hz and the amplitude is 2 mm. Steel balls having a diameter of 10 mm, made of SUJ-2, were used as the opposite material. The lubricant was not used. The case where the load of 5000 N is applied is shown by symbol ◎. The case where the load of 4000 to 5000 N or

more is applied is shown by symbol ○. The case where the load of 2000 to 4000 N is applied is shown by symbol △. The case where the load of less than 2000 N is applied is shown by symbol ×.

5 Porosity:

Porosities were measured in conformity with ISO2738: Permeable sintered metal materials determination of densit B, oil content and open porosit B.

Pore size ratio:

10 Pore size ratios were obtained under a microscope and using a pore quantity device. First, pores having an inlet diameter of 10 to 200 μm were obtained. Next, the corresponding inner pore diameters were obtained. Average values were obtained by calculating quotients.

15 The above-mentioned result proves that the sintered material of the present invention excels in the adhesive property of the film, which is formed on the sintered material through the selective chemical etching solution process. Furthermore, the sintered material has a good
20 corrosion resistance and a good sliding property. That is, even in the case of the comparative example 1 that carries out the simple etching only but does not carry out the selective chemical etching, a skin is formed on the sintered material, in a manner similar to those in the
25 embodiments. However, the skin is poor in adhesive

property, corrosion resistance, and sliding property.

According to the present invention, a film of a solid lubricating agent such as PTFE or molybdenum disulfide can be fixed firmly and in adhesive state on the sintered
5 material. Good sliding properties can be obtained under unlubricating conditions or under corrosive conditions.